

Figure 1

Minimum signal peptide score	false positive rate	false negative rate	proba(0.1)	proba(0.2)
3,5	0,121	0,036	0,467	0,664
4	0,096	0,06	0,519	0,708
4,5	0,078	0,079	0,565	0,745
5	0,062	0,098	0,615	0,782
5,5	0,05	0,127	0,659	0,813
6	0,04	0,163	0,694	0,836
6,5	0,033	0,202	0,725	0,855
7	0,025	0,248	0,763	0,878
7,5	0,021	0,304	0,78	0,889
8	0,015	0,368	0,816	0,909
8,5	0,012	0,418	0,836	0,92
9	0,009	0,512	0,856	0,93
9,5	0,007	0,581	0,863	0,934
10	0,006	0,679	0,835	0,919

Figure 2

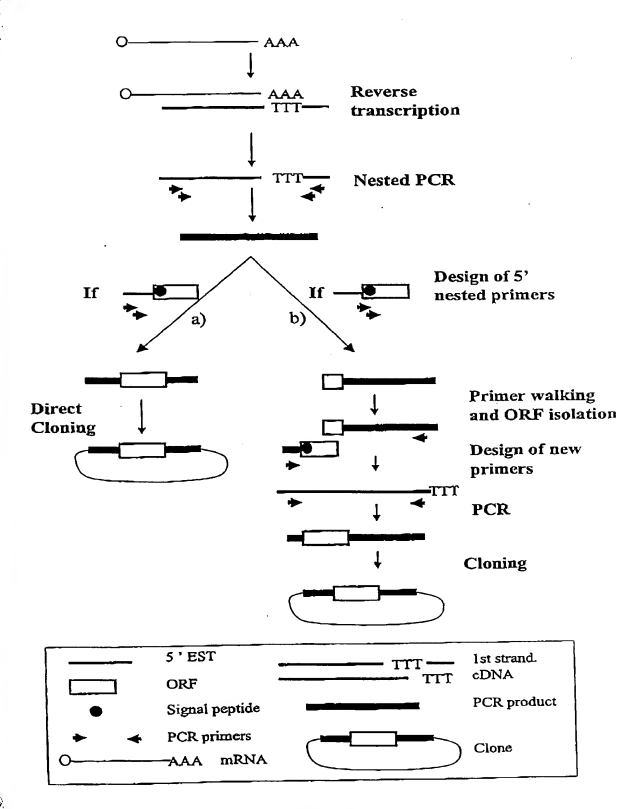


Figure 3

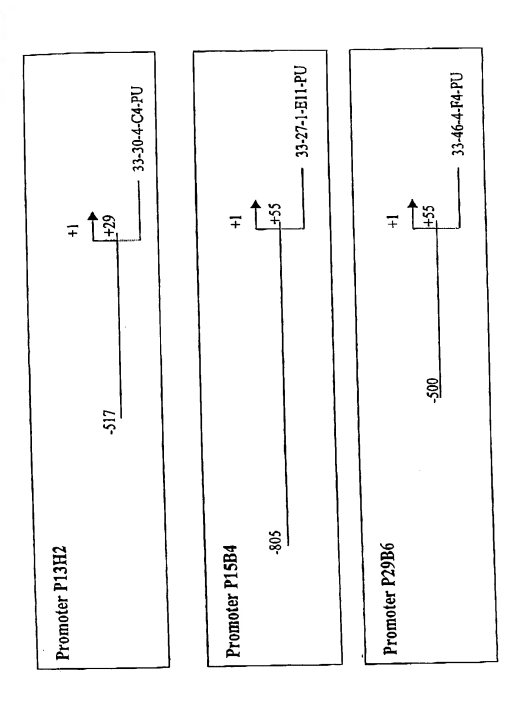


Figure 4

Promoter sequence P13H2 (546 bp):

		Orient			
Matrix	Position	ation	Scor	Length	Sequ nce
CMYB_01	-502	+	0.983	9	TGTCAGTTG
MYOD_Q6	-501	_	0.961	10	CCCAACTGAC
S8_01	-444	-	0.960	11	AATAGAATTAG
S8 01	-425	+	0.966	11	AACTAAATTAG
DELTAEF1_01	-390	-	0.960	11	GCACACCTCAG
GATA_C	-364	_	0.964	11	AGATAAATCCA
CMYB_01	-349	+	0.958	9	CTTCAGTTG
	-343	+	0.959	14	TTGTAGATAGGACA
GATA1_02	-339	+	0.953	11	AGATAGGACAT
GATA_C	-235	+	0.973	16	CATAACAGATGGTAAG
TALIALPHAE47_01	-235	+	0.983	16	CATAACAGATGGTAAG
TAL1BETAE47_01	-235	+	0.978	16	CATAACAGATGGTAAG
TAL1BETAITF2_01	-232	·	0.954	10	ACCATCTGTT
MYOD_Q6		-	0.953	13	TCAAGATAAAGTA
GATA1_04	-217		0.963		AGTTGGGAATTCC
IK1_01	-126		0.985		AGTTGGGAATTC
IK2_01	-126		0.962		TGGGAATTCC
CREL_01	-123				TCAGTGATATGGCA
GATA1_02	-96		0.950		TAAAACAAAACA
SRY_02	-41		0.951	_	TITAGCGC
E2F_02	- 3 3		0.957	_	TGAGGGGA
MZF1_01	-5	-	0.975	. 6	(0/100001

Promoter sequence P15B4 (861bp):

		Oneni			
Matrix	Position	ation	Score	Length	Sequence
	-748	-	0.956	11	GGACCAATCAT
NFY_Q6	-738	+	0.962	8	CCTGGGGA
MZF1_01	-684	+	0.994	9	TGACCGTTG
CMYB_01			0.985	9	TCCAACGGT
VMYB_02	-682	_	0.968	9	TTCCTGGAA
STAT_01	-673	+	0.951	9	TTCCAGGAA
STAT_01	-673	-		8	TTGGGGGA
MZF1_01	-55 6		0.956		GAATGGGATTTC
IK2_01	-4 51	+	0.965	12	AGAGGGGA
MZF1_01	-424	+	0.986	8	
SRY 02	-398	-	0.955	12	GAAAACAAAACA
MZF1 01	-216	+	0.960	8	GAAGGGGA
MYOD_Q6	-190	+	0.981	10	AGCATCTGCC
	-176		0.958	11	TCCCACCTTCC
DELTAEF1_01	5		0.992	11	GAGGCAATTAT
S8_01	16		0.986	_	AGAGGGGA
MZF1_01	10	, -	5.000		

Promoter sequence P29B6 (555 bp):

	C	Drient			
Matrix	Position (ation	Score	Length	Sequence
•••	-311	+	0.964	16	GGACTCACGTGCTGCT
ARNT_01	-309	+	0.965	12	ACTCACGTGCTG
NMYC_01	-309	+	0.985	12	ACTCACGTGCTG
USF_01	-309	_	0.985	12	CAGCACGTGAGT
USF_01	-309 -309	_	0.956	12	CAGCACGTGAGT
NMYC_01	- - ·		0.972	12	CAGCACGTGAGT
MYCMAX_02	-309	•	0.997	8	TCACGTGC
USF_C	-307	+		8	GCACGTGA
USF_C	-307	-	0.991	8	CATGGGGA
MZF1_01	-292	-	0.968	_	CTCTCCGGAAGCCT
ELK1 02	-105	+	0.963	14	TCCGGAAGCC
CETS1P54_01	-102	+	0.974	10	AGTGACTGAAC
AP1_Q4	-42	-	0.963	11	AGTGACTGAAC
AP1FJ_Q2	-42	-	0.961	11	TGTGGTCTC
PADS C	45	+	1.000	9	101001010

Figure 5

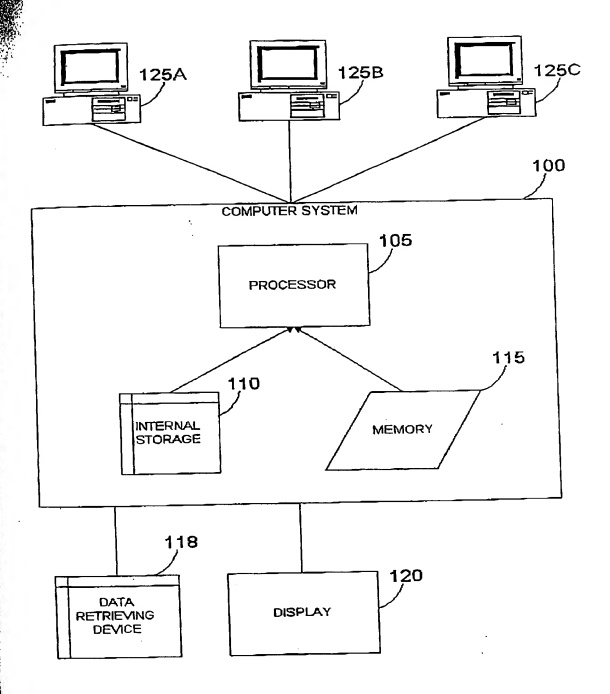


FIGURE 6

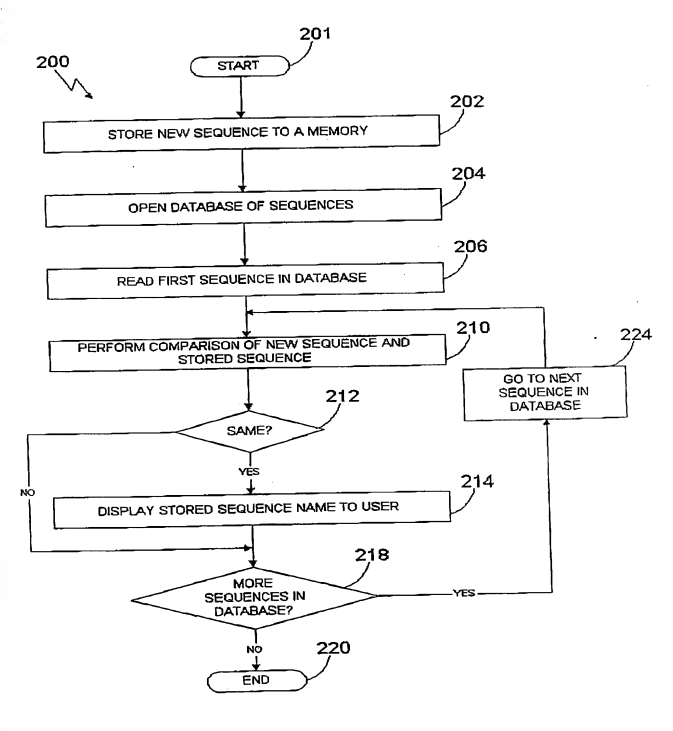
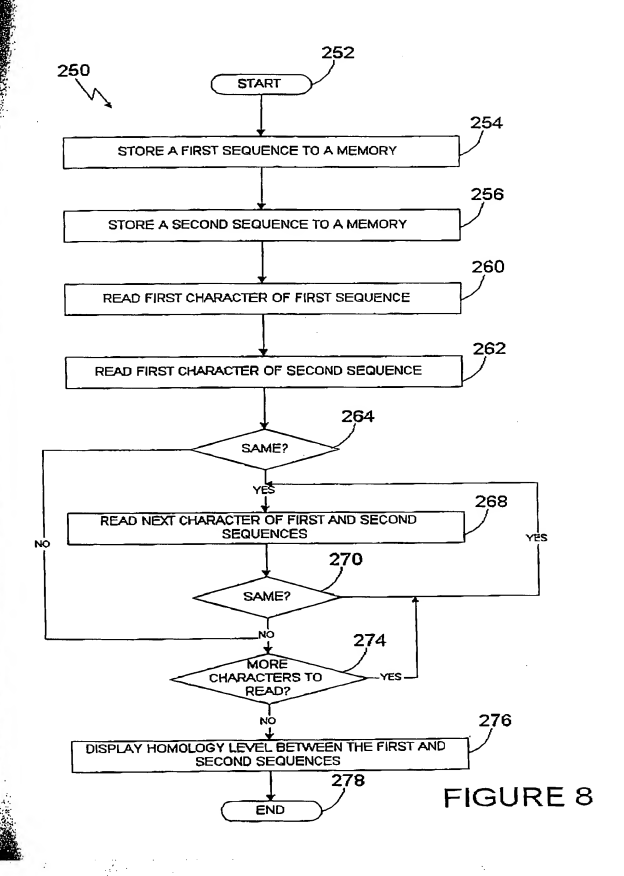
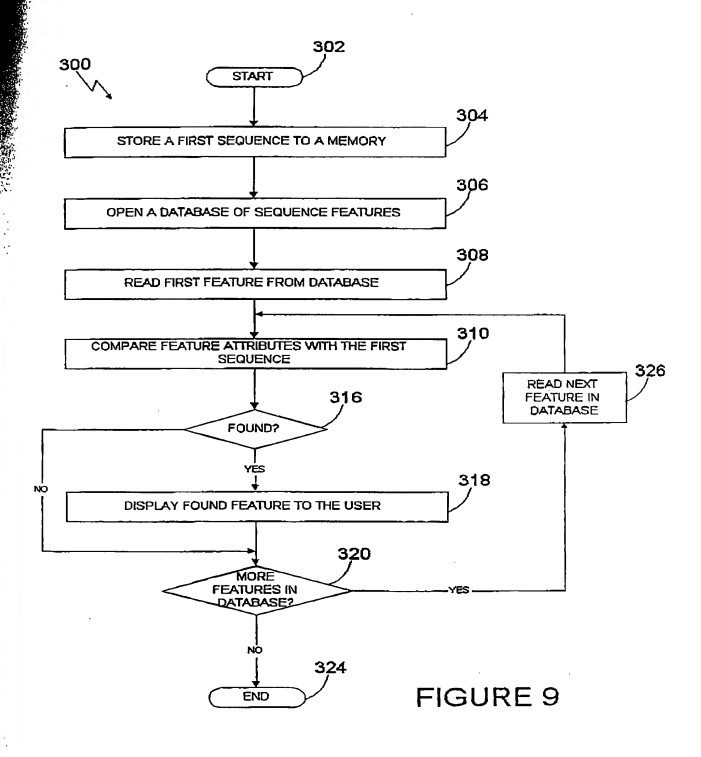


FIGURE 7





	Search characteristic	istic			Selection	Selection Characteristics
Step	Program	Strand	Parameters	Identity (%) Length (bp)	Length (bp)	Comments
miscellanaeous	FASTA	both	•	06	51	
tRNA	FASTA	both	•	08	09	
rrna	BLASTN	both	S=108	08	8	
mtRNA	BLASTN	both	S=108	08	40	
Procaryotic	BLASTN	both	S=144	06	40	
Fungal	BLASTN	both	S=144	96	8	
Alu	BLASTN	both	S=72, B=5	0/	40	max 5 matches, masking
L1	BLASTN	both	S=72, B=5	0%	40	max 5 matches, masking
Repeats	BLASTN	poth	S=72	01	40	masking
			W=6, S=10, E=1000,			
PolyA	BLAST2N	top	N=12	96	01	in the last 100 nucleotides
Polyadenylation signal	•	dog	AATAAA all	AATAAA allowing 1 mismatch	atch	in the 50 nucleotides before the 5' end of the polA
Vertebrate	BLASTN then FASTA	poth	•	90 then 70	30	first BLASTN, then FASTA on maching sequences
ESTs	BLAST2N	both	-	06	30	
Geneseq	BLASTN	both	W=8, B=10	06	30	
ORF	BLASTP	top	W=8, B=10		•	on ORF proteins, max 10 matches
Proteins	BLASTX	top	E = 0.001	202	30	

1

Figure 10